## PATENT SPECIFICATION

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## (54) STABILIZED TRANSPARENT RECEPTOR SHEET

(71) We, MINNESOTA MINING AND MANUFACTURING COMPANY, a corporation organized and existing under the laws of the State of Delaware, United States of America, of 3M Center, Saint Paul, Minnesota 55101, United States of America, do hereby declare the invention for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement: -

This invention relates to transparent receptor sheets. Particularly, the invention relates to the stabilization of receptor sheets capable of being imaged containing reducible metal soap and organic reducing agent.

In United States Patent No. 3,935,012, there are described light-sensitive intermediate sheets comprising (a) photolyzable heat-stable source of hydrogen halide, (b) acid-cleavable adduct of a first reactant and an organic compound containing a vinyl ether linkage, the adduct being capable of supplying a reducing agent when cleaved with acid, and (c) acid acceptor for the hydrogen halide in an amount less than sufficient to retain all of the hydrogen halide available from the photolyzable source upon exposure to a light image. When this sheet is exposed to a light image hydrogen halide is formed which cleaves the adduct so as to free a reducing agent at light-exposed areas. The intermediate sheet is then placed against a receptor sheet containing a second reactant (e.g., reducible metal soap and organic reducing agent such as sterically hindered phenol) and heated so as to induce a visible image-forming reaction. An image is formed in the receptor sheet due to reduction of a portion of the metal soap (to the metal) by reaction thereof with the liberated reducing agent from the light-struck areas of the intermediate sheet. This then caralyzes the heatinduced reduction of the remaining metal soap by the less reactive organic reducing agent originally present in the receptor to form a visible image.

Although the receptor sheet which has been imaged in accordance with the foregoing procedure is of good quality and is very easy to use, it has been found that those receptor sheets which include reducible metal soap and organic reducing agent tend to exhibit unacceptable increases in background optical density when exposed for a prolonged period of time (e.g. 2 hours or more) on a conventional overhead projector. During image formation, a small amount of material from the non-light-exposed areas of the intermediate sheet is transferred to the receptor sheet. Thus, when the imaged receptor sheet is used on an overhead projector, sufficient reducing agent is liberated so as to bring about the increase in background optical density. We have sought to overcome this problem of background optical density increase by including in the receptor sheet a small amount of aromatic dye.

Accordingly the present invention provides a receptor sheet for use in an image transfer process comprising a transparent support having at least one coating layer thereover including a film-forming binder, a reducible metal soap, an organic reducing agent and from 0.00007 to 0.005 parts by weight, per part of reducible metal soap, of a polycyclic aromatic dye compound exhibiting a development characteristic, as herein defined, of at least 150 seconds.

Optionally, the coating may also contain a toner (e.g. phthalazinone) for the metal image, or fillers or various other additives.

Preferably the support is a thin, flexible, transparent plastics film of a type which is known in the art as being useful for use in a projection transparency.

The improved receptor sheets of the present invention, after being imaged, may be projected for prolonged periods on conventional overhead projectors without objec-

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	tionable increase in al	2
	the polycyclic aromatic dye stabilizes the receptor sheet by retarding the reduction of The reducible metal soap by the liberated reducing agent in background areas.	
	the metal soap by the liberated make the receptor sheet by retarding the reduction at	
5	the metal soap by the liberated reducing agent in background areas.  The reducible metal soap present in the receptor sheet is preferably a silver soap metal soaps such as ferric stearing, silver stearate), although other metal soaps such as ferric stearing the reducible metal soaps.	
,		
	metal soaps such as ferric stearate, gold stearate and cerium stearate are also useful.  The organic reducing search known in the art.	
	Other useful reducible metal soaps are known in the art.  The organic reducing agents which	. 5
	The organic reducing agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents agents which are useful in the receptor sheets are also bis - pyrogalol: 4 - steered agents agent agents ag	
10	well known in the art and include, as representative examples, pyrogalol; 4 - azeloyl - gallic acid anilide; methyl gyllagallol; gallacetophenone; diteriaryhygal	
10	bis - pyrogallol; 4 - stearoyl pyrogallol; gallacetophenone; ditertiarybutyl pyrogallol; gallate; dodecyl gallate, gallate, ethyl gallate; normal and isopropyl gallate;	
	gallic acid anilide; methyl gallate, ethyl gallate; normal and isopropyl gallate; butyl protocatechuate; 2.5-dibyd-methyl gallate; ammonium gallate; ethyl protocatechuate; 2.5-dibyd-methyl gallate; butyl ga	
	galate; dodecyl gallate, gallic acid: amponium ormal and isopropyl gallate; butyl	-10
	gallate; dodecyl gallate, gallic acid; ammonium gallate; ethyl protocatechuate; butyl protocatechuate; 2,5-dihydroxy benzoic acid; 1-hydroxy-2-naphthoic acid; 2-hydroxy, sodium gallate; protocatechuate; catechol; 2,3-naphthalene diol; 4 layers,	
15	3-naphthoic acid; phloroglucinol; catechol; 2,3-naphthalene diol; 2-hydroxy, sodium gallate; protocatechualdehyde; 4-methyl esculetin; 3,4-dihydroxy benzoic acid; 1-hydroxy benzoic acid; 2-hydroxy, 2,3-dihydroxy benzoic acid; 1-hydroxy benzoic ac	
	sodium gallate; protocatechualdehyde; 4-methyl esculetin; 3,4-dihydroxy benzoic acid; hydroquinone; 4,4'-dihydroxy benzoic acid; hydroquinone; 4,4'-dihydroxy benzoic acid;	
	phenylorety benzoic acid; hydroquinone: 4/ dibud; 3,4-dihydroxy benzoic acid;	1.0
	2,3-dihydroxy benzoic acid; hydroquinone; 4,4'-dihydroxy biphenyl; 3,4-dihydroxy benzoic acid; phenylacetic acid; 4(3',4'-dihydroxyphenylazo) benzoic acid; 4(3',4'-dihydroxyphenylazo) benzoic acid; 2,2-methylazo	15
	phenylacetic acid; hydroquinone; 4,4'-dihydroxy biphenyl; 3,4-dihydroxy-trihydroxybenzoic acid; ortho- and paraphenylene diamine; tetramethyl benzidine; beta naphthols; 4-methylary 1-a-tylene phenylmethane; o-, m- and p-aminobenzoic acid; ortho- beta naphthols; 4-methylary 1-a-tylene	
20	4,4',4"-diethylamino triphenylmethane; o-, m- and p-aminobenzoic acid; 2,2-methylene bis-3,4,5-beta naphthols; 4-methoxy-1-naphthol, 1-hydroxy-dihydronaphthalene; o-, m- and hydroquinoline. These compounds in the compound in the compounds in the compound in the compounds in the compound in the compounds in the compound in th	
	hydroguipoline 7-methoxy-1-naphthol, 1-hydrogu dibadonic acids; alpha and	
	hydrogen atom areast compounds are cyclic or aromatic compounder, and tetra-	20
	is attached to an atom of carbon, overgen an active	20
	of metal ion and atom of the cyclic ring. They are capable of nitrogen which in turn	
25	is attached to an atom of carbon, oxygen or nitrogen which in turn of metal ion and precipitation of metal on being dissolved at moderate sequences.	
	The reducible metal sait in an organic solvent	
	prior to coating but said organic reducing agent	25
	the aromatic dye being present in the layer continuing the reducible metal salt.	23
	Various trace of Silver in the layer containing the red with	
30	example one may be used in the last sail.	
	acetate, polyvinyl acceptors propionate resin, polyethyl method icceptor sheets. For	
	Other conventional bind chloride copolymer, and called resin, polyvinyl	30
	The relative conditions are well known in the art	
	in the recentor many of reducible metal soan and ormalisment	
35	so as to be capable of forming a visible and distinct image when heated in the present of the reactant (i.e., reducing agent) supplied by the intermediate sheet	
	of the reaction (in the reaction of the reaction of the present	
	of the reactant (i.e., reducing agent) supplied by the intermediate sheet described parts by weight per part of organic reducing agent present.  The types of aromatic divariations and distinct image when heated in the presence above, Generally speaking, the amount of reducible metal soap present is from 4 to 12.	35
	parts by weight per part of organic reducible metal soap present is from 4 to 12.  The types of aromatic dyes which are metal present.	
40	The types of aromatic does which reducing agent present.	
40	MULCASE in bookers	
	of chemical classification of dyestiffs. Fig. 1 and been entirely definable in the same	
	of chemical classification of dyestuffs. Fortunately, however, a simple test has been useful and those which are not. This test is used herein to define the double are characteristics. This test procedure.	40
	useful and those which are not. This test is used herein to define the development the following ingredients in the arm as follows: A solution is first procedure to the following ingredients in the arm as follows:	
45	tharacteristics. This test procedure is as fall used herein to define the development	
43	characteristics. This test procedure is as follows: A solution is first prepared using	
	C the control of the	
		4 -
	Cellulose acetate propionate ("FAP	45
	Cellulose acetate propionate ("EAP 482—20"), available from Fastman	45
	Methylethylethone from Eastman) 5 grams	45
o	Methylethylketone S grams Aromatic due to be a promise of the second sec	45
o	Methylethylketone 5 grams Aromatic dye to be tested 95 grams 4-methoxy-1-paphthol 0.10 millimoles	45
o	Methylethylketone Aromatic dye to be iested 4-methoxy-1-naphthol  Then 15 grams 6 to 10 millimoles  0.575 millimoles	
0	Methylethylketone Aromatic dye to be iested 4-methoxy-1-naphthol  Then 15 grams of this solution is poured into an element of 0.5 inch) and involved into 0.5 inch in our element of 0.5 inch in ou	45 50
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5	Methylethylketone Aromatic dye to be tested 4-methoxy-1-naphthol  Then 15 grams of this solution is poured into an aluminium dish (to a depth General Electric tungsten filament light source. Thereafter the irradiated solution is and which is described below), followed by drying at 6500.	
5	Methylethylketone Aromatic dye to be tested 4-methoxy-1-naphthol  Then 15 grams of this solution is poured into an aluminium dish (to a depth General Electric tungsten filament light source. Thereafter the irradiated solution is and which is described below), followed by drying at 65. After drying, the receptor receptor sheet at 100°C. (212°F.). The time required for the standard of the standard for the receptor sheet at 100°C. (212°F.).	50
5	Methylethylketone Aromatic dye to be tested 4-methoxy-1-naphthol  Then 15 grams of this solution is poured into an aluminium dish (to a depth General Electric tungsten filament light source. Thereafter the irradiated solution is and which is described below), followed by drying at 65°C. After drying, the receptor sheet to show an increase to 0.85 is recorded to the optical density of the sistic, for the partial.	
5	Methylethylketone Aromatic dye to be iested 4-methoxy-1-naphthol  Then 15 grams of this solution is poured into an aluminium dish (to a depth of 0.5 inch) and irradiated at a distance of 6 inches for 1 hour with a 150 watt coated (at 50 microns wet thickness) onto a receptor sheet (6 months or less in age sheet is heated at 100°C. (212°F.). The time required for the optical density of the invention or his in the particular dye being tested. Those dates which is development character-invention or his in the standard of the standard of the invention or his in the standard of the standard of the invention or his in the standard of the standard of the invention or his in the standard of the standard of the standard of the invention or his in the standard of th	50
5	Methylethylketone Aromatic dye to be iested 4-methoxy-1-naphthol  Then 15 grams of this solution is poured into an aluminium dish (to a depth of 0.5 inch) and irradiated at a distance of 6 inches for 1 hour with a 150 watt coated (at 50 microns wet thickness) onto a receptor sheet (6 months or less in age sheet is heated at 100°C. (212°F.). The time required for the optical density of the invention exhibit a "development characteristic" of at less 150 which are useful in the present dyes exhibit a "development characteristic" of at less 150 grams  95 grams  0.10 millimoles 0.575 millimoles 150 watt	50
5	Methylethylketone Aromatic dye to be iested 4-methoxy-1-naphthol  Then 15 grams of this solution is poured into an aluminium dish (to a depth of 0.5 inch) and irradiated at a distance of 6 inches for 1 hour with a 150 watt coated (at 50 microns wet thickness) onto a receptor sheet (6 months or less in age sheet is heated at 100°C. (212°F.). The time required for the optical density of the invention exhibit a "development characteristic" of at less 150 which are useful in the present dyes exhibit a "development characteristic" of at less 150 grams  95 grams  0.10 millimoles 0.575 millimoles 150 watt	50
5	Methylethylketone Aromatic dye to be iested 4-methoxy-1-naphthol  Then 15 grams of this solution is poured into an aluminium dish (to a depth of 0.5 inch) and irradiated at a distance of 6 inches for 1 hour with a 150 watt coated (at 50 microns wet thickness) onto a receptor sheet (6 months or less in age sheet is heated at 100°C. (212°F.). The time required for the optical density of the istic" for the particular dye being tested. Those dyes which are useful in the present dyes exhibit a "development characteristic" of at least 150 seconds. Preferably the procedure the optical density is measured using a conversion.	50 55
5	Methylethylketone Aromatic dye to be iested 4-methoxy-1-naphthol  Then 15 grams of this solution is poured into an aluminium dish (to a depth of 0.5 inch) and irradiated at a distance of 6 inches for 1 hour with a 150 watt coated (at 50 microns wet thickness) onto a receptor sheet (6 months or less in age sheet is heated at 100°C. (212°F.). The time required for the optical density of the istic" for the particular dye being tested. Those dyes which are useful in the present dyes exhibit a "development characteristic" of at least 150 seconds. Preferably the procedure the optical density is measured using a conversion.	50
5	Methylethylketone Aromatic dye to be iested 4-methoxy-1-naphthol  Then 15 grams of this solution is poured into an aluminium dish (to a depth of 0.5 inch) and irradiated at a distance of 6 inches for 1 hour with a 150 watt coated (at 50 microns wet thickness) onto a receptor sheet (6 months or less in age sheet is heated at 100°C. (212°F.). The time required for the optical density of the invention or his in the particular dye being tested. Those dates which is development character-invention or his in the standard of the standard of the invention or his in the standard of the standard of the invention or his in the standard of the standard of the invention or his in the standard of the standard of the standard of the invention or his in the standard of th	50 55

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	dispersion	is	prepared	using	the	following	ingredients	in	the	amounts
shown:										

	Parts by Wt.
Silver behenate Tetrachlorophthalic anhydride Stannous stearate Methylethylketone	12.00 0.24 0.0072 87.7528
	100.0000

The above ingredients are mixed for one-half hour, then homogenized at 8000 psi, cooled to 80°F., and homogenized again at 8000 psi.

A binder solution is prepared with the following ingredients:

		Parts	
15	Tenite H4® cellulose acetate butyrate resin (available from Eastman) Elvacite 2042® polyethylmethacrylate	7.50	15
	resin (available from E. I. du Pont)	7.50	13
	Methylethylketone	85.00	
		100.00	

A coating composition is then prepared using the following ingredients:

20		Parts	20
	Silver behenate dispersion	5 <b>4.5531</b>	
	Binder solution	43.6400	
	Phthalazinone	0.9819	
	2,6-di-t-butyl-p-cresol	0.7850	
25	4,4'-methylene-bis-2,6-di-t-butylphenol	0.0400	25
		100.0000	

The above ingredients are thoroughly blended and then coated onto a transparent plastics film at an orifice of 3.5 mils (87.5 microns) and dried in a forced air oven (3 minutes at 180°F.) to leave a dried, clear and transparent coating of 0.70 grams per square foot. A top coating is then applied using a 5% solution of cellulose acetate propienate in methylethylketone (coated at a wet thickness of 50 microns and dried to leave a dry coating having a thickness of 5 microns).

A preferred class of useful polycyclic aromatic dyes having a "development characteristic" of at least 150 seconds are oxygen-containing dyes comprising at least two moieties in conjugate relationship as part of a single chromophore, each moiety comprising three linearly kata condensed six-membered aromatic rings, an —OZ group being attached to the meso position of each moiety, wherein Z is a stable, monovalent radical; the —OZ group being a solubilizing group for the compound. Preferably, at least one auxochromic group comprising an atom having an atomic weight of at least 31 is bonded to the chromophore, the atom being attached directly to the chromophore. The above-mentioned moieties comprising three linearly kata condensed six-membered aromatic rings may be described pictorially as follows:

The aromatic rings may be homocyclic (carbon atoms) or heterocyclic, the hetero atoms being generally nitrogen. Of course, a single compound can contain both homocyclic and heterocyclic moieties of the type just described, or the compound may contain only homocyclic or only heterocyclic moieties.

These moieties are in conjugate relationship in the dye compound so that these moieties are part of a single chromophore (i.e., part of the same chromophore). A

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chromophore may be defined as a group of atoms or electrons in a molecule which is chiefly responsible for an absorption band, as defined in Theory and Application of Ultraviolet Spectroscopy; Jaffe and Orchin; John Wiley & Sons, Inc., (1962). The two moieties may be bonded or condensed together in peri fashion, e.g.,

wherein the two moieties share atoms, or the moieties may be attached to each other in conjugate relationship through at least one linking moiety. The linking moieties are selected from (a) atoms which are at least trivalent and which are capable of forming covalent bonds, e.g., nitrogen, carbon, sulphur and oxygen, (b) ligands, having two or more atoms, which are at least bidentate (e.g., such ligands can be bidentate, tridentate, quadradentate, hexadentate, octadentate, etc.), and (c) covalent

The size, chemical nature or structure of the linking moiety is not critical so long as the two moieties comprising the kata condensed aromatic rings are attached to each other in conjugate relationship such that they are part of a single

The linking moiety, of course, can be a polycyclic structure (homocyclic or heterocyclic). Hetero atoms in the heterocyclic linking moieties are generally nitrogen,

In many of the compounds there are two linking moieties. For example, there may be two ligands, one ligand and a covalent bond or two covalent bonds as linking

Preferably, these dye compounds have ar least one auxochromic group bonded to the chromophore thereof. These preferred dyes are described in U.S. 3,819,664. The auxochromic groups may consist of one atom or of many atoms, so long as a heavy atom (i.e., having an atomic weight of at least 31), present as part of the auxochromic group, is bonded directly to the chromophore portion of the dye. An auxochromic group may be defined as a group bonded to a chromophore which influences the nature of the excited states, as defined in Theory and Application of Ultraviolet Spectroscopy; Jaffe and Orchin; John Wiley & Sons, Inc. (1962).

Preferred single atom auxochromic groups include chlorine, bromine, mercury, sulphur, iodine and selenium. Other useful single atom auxochromic groups include phosphorus (treated berein as basic as basic as basic as a supply supply as a supply as a supply suppl phosphorus (treated herein as having an atomic weight of 31), arsenic, tellurium, germanium, tin, lead and antimony. The auxochromic group may consist of more than one atom so long as a heavy atom present as part of the auxochromic group is

directly bonded to the chromophore portion of the dye.

Although the —OZ and —OZ' groups which are present on the dye compounds are also auxochromic groups, it has been found that the —OZ and —OZ' groups are also auxochromic groups, it has been found that the —OZ and —OZ' groups are primarily solubilizing groups for the compound in their primarily determines. are primarily solubilizing groups for the compound, i.e., they primarily determine the solubility of the dye compound in various solvent media into which the dye may be placed. Thus, the chemical structure and nature of the Z and Z' radicals are not

Generally, it may be said that Z and Z' are monovalent radicals which are stable under ambient conditions and which do not cause decomposition of the chromophore portion of the dye compound. That is, these radicals do not oxidize or reduce the chromophore portion of the dye compound nor do they destroy or adversely affect the effectiveness of the compound as a dye. Within these limitations the —OZ and —OZ' radicals can be stable derivatives of an inorganic acid, e.g., —OSO<sub>3</sub>Y+ where Y is an alkali metal, alkaline earth metal, or ammonium ion; —OPO<sub>3</sub>R<sub>2</sub> where R is hydrogen or a stable organic radical; and —OPO<sub>2</sub>R<sub>2</sub> where R is hydrogen or a

Insofar as organic radicals are concerned, Z and Z' may be alkyl, cycloalkyl, substituted alkyl and cycloalkyl, alkenyl, alkynyl, aryl, polycyclic, acyl, alkaryl or aralkyl. Z and Z' may be the same or different. Alkyl radicals having one carbon or 10

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more are common Z radicals, and lower alkyl radicals containing from 1 to 6 carbon atoms are preferred, although long chain alkyls are also useful. Substituted alkyl radicals are herein defined to include alkyl radicals which are substituted with any moiety or group other than hydrogen atoms and other alkyl radicals.

In addition to the —OZ and —OZ' solubilizing groups and the auxochromic groups the above classes of dye compounds may also be substituted with various other groups (e.g., fluorine, nitrile, hydroxy, alkyl, aryl, polycyclic, acyl, alkoxy) which are stable and do not cause decomposition of the chromophore portion of the dye compound.

The dye compounds used in this invention can be prepared by independent synthesis but they are more conveniently prepared from precursor dyes. For example, Vat Violet 7 (Colour Index No. 59321) may be alkylated according to the following scheme with conventional alkylating agents:

where RX represents agents such as alkyl iodide, alkyl tosylate, alkyl benzene sulphonate, and dialkyl sulphate. Other representative useful alkylated vat dyes include ethylated Vat Blue 18 (Colour Index No. 59815), and ethylated Vat Green 1 (Colour Index No. 59825).

To obtain compounds having R groups which are allyl, substituted allyl, propargyl, or substituted propargyl, alkylating agents such as allyl bromide and propargyl bromide may be used in the above reaction scheme.

To obtain compounds wherein R and R' represent cycloalkyl radicals the dianion may be reacted with acrivated cycloalkyl halides, such as 2-chlorocyclohexanone, using the above reaction scheme. The resulting product has the following formula:

This product may be subsequently reduced via a Wolff-Kischner type reaction to yield a compound of the following formula:

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50 The above ingredients were thoroughly blended and then coated onto a trans-100.0000 parent plastics film at an orifice of 6 mils (150 microns) and dried in a forced air oven (3 minutes at 180°F.) to leave a dried, clear and transparent coating of 0.0013 50

Butylated Vat Violet 7 (Colour Index

No. 59321) dye

A top coating was then applied using a 5% solution of cellulose acetate propionate

0.9819

0.7850

0.0080

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1,546,223 7 in methylethylketone (coated at a wet thickness of 50 microns and dried to leave a dry coating having a thickness of 5 microns). Example 2 An intermediate sheet was prepared using a coating composition comprising: 5 **Parts** 5 Cellulose acetate butyrate 7.887 Solvent (methylethylketone) 91.00 Adduct of 4-methoxy-1-naphthel and dihydropyran (equimolar) 0.717 10 Tris-(tribromomethyl)-s-triazine 0.58 10 Spectral sensitizer (perylene) 0.052 Acid acceptor (diphenylguanidine) 0.016 The above composition was coated onto a clear polyester film from a 3 mil (75 microns) coating orifice and dried, all operations being conducted under appropriate safe light, to provide a photosensitive intermediate sheet. 15 15 The coated surface of the resulting intermediate sheet was then placed in contact with a printed original having light-absorptive image areas on a reflective white background, followed by uniformly exposing through the intermediate (i.e., in reflex position) to intense illumination from a bank of tungsten filament lamps for a time just sufficient to develop the coating completely at the background areas. A typical exposure time is 12 to 15 seconds. The image areas were also affected but to a signifi-20 20 cantly lesser extent due to absorption in the image areas of the original of a portion of the illumination. The exposed intermediate sheet was then placed with its coated surface against the coated face of the receptor sheet of Example 1 (e.g., between rolls or platens) and heated for 4 to 5 seconds at 125 to 140°C. A sharp negative copy of 25 25 the graphic original was obtained on the receptor sheet. The receptor sheet was then placed on a "Model AR-66" overhead projector (available from Minnesota Mining and Manufacturing Company) for 16 hours with no perceptible increase in background optical density. This result was distinguished from that observed when the example was repeated 30 30 using a receptor sheet having no polycyclic aromatic dye therein. In this latter case there was a 1000% increase in background optical density when left on the above projector for 16 hours. Examples 3 to 7 Several receptor sheets were prepared, each having a different amount of poly-35 35 cyclic aromatic dye therein. A stock solution was first prepared using the following ingredients: **Parts** Silver behenate dispersion from Example 1 54.585 40 Phthalazinone 0.982 40 Binder solution from Example 1 43.585 2,6-di-t-butyl-p-cresol (sterically hindered phenol reducing agent) 0.793 To similar samples of the above stock solution (each sample containing 25 grams 45 of silver behenate soap) were added varying amounts of butylated Vat Violet 7 (poly-45 cyclic aromatic dye), as shown in Table I: TABLE I Polycyclic Example No. Dye Added (Grams) 50 3 (Comparative) 50 0.002 0.004 6 7 (Comparative) 0.008 0.016 55

Each solution was then knife coated onto separate samples of 2 mil (50 microns)

thick polyester film using a 100 micron orifice. The samples were then air dried at 190°F. (88°C.) to leave a dried coating 20 microns thick, after which a protective top

coating of cellulose acetate butyrate was applied (5 microns, dry thickness).

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projector for a prolonged period and the background optical density was measured at various times. The results are shown in Table IV.

## TABLE IV

			Background	Optical Density
	Example No	o. Initial	4 Hours	12 Hours
	<b>i</b> 4	0.06	0.14	1.15
	15	0.06	0.07	0.09

Other variants are possible within the scope of the present invention.

## WHAT WE CLAIM IS:-

1. A receptor sheet for use in an image transfer process comprising a transparent support having at least one coating layer thereover including a film-forming binder, a reducible metal soap, an organic reducing agent and from 0.00007 to 0.005 parts by weight, per part of reducible metal soap of a polycyclic aromatic dye compound exhibiting a development characteristic, as herein defined, of at least 150 seconds.

exhibiting a development characteristic, as herein defined, of at least 150 seconds.

2. A receptor sheet as claimed in Claim 1, wherein the dye compound comprises at least two moieties in conjugate relationship as part of a single chromophore, each moiety comprising three linearly kata condensed six-membered aromatic rings having an —OZ group attached to the meso position of each moiety, wherein Z is a stable, monovalent radical and wherein the —OZ group is a solubilizing group for the dye compound.

3. A receptor sheet as claimed in Claim 1 or 2, wherein the dye compound is of the formula

where R and R' are each an alkyl group containing from 1 to 6 carbon atoms.

4. A receptor sheet as claimed in Claim 3, wherein R is a butyl group.

5. A receptor as claimed in claim, wherein the dye compound is octaphenyl-

tetrazaporphyrin or erythrosin.

6. A receptor sheet as claimed in any of the preceding claims, wherein the reducible metal soap is silver behenate, silver stearate, ferric stearate, gold stearate or cerium stearate.

7. A receptor sheet as claimed in any one of the preceding claims, wherein the organic reducing agent is a sterically hindered phenol.

8. A receptor sheet as claimed in Claim 7, wherein the organic reducing agent is 2,6-di-t-butyl-p-cresol, or 4,4'-methylene-bis-2,6-di-t-butylphenol.

9. A receptor sheet as claimed in any one of Claims 1 to 6, wherein the organic reducing agent is 4-methoxy-1-naphthol.

10. A receptor sheet as claimed in any one of the preceding claims, wherein the dye compound is ethylated Vat Blue 18 (Colour Index No. 59815).

11. A receptor sheet in accordance with Claim 1 and substantially as herein described with reference to the Examples.

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